## PATENT SPECIFICATION

(11)1 400 898.

5

10

15

20

25

30

35

(21) Application No. 33482/73 (22) Filed 13 July 1973

(31) Convention Application No. 271 943 (32) Filed 14 July 1972 in (33) United States of America (US)

(44) Complete Specification published 23 July 1975

(51) INT CL\* C11D 3/60; D06M 11/06//(ClifD 3/60, 1/02, 1/88, 3/02, 3/12)

(52) Index at acceptance

C5D 6A2 6A5A 6A5B 6A5C 6A5D1 6A5D2 6A5F 6A9 6B12A 6B12B1 6B12F1 6B12F2 6B12G1 6B12G2A 6B12G4 6B1 6B2 6C6

DIP 1A3 1A5 A18 B6 CIX

(72) Inventors THOMAS D. STORM and IOSEPH P. NIRSCHL.



## (54) DETERGENT COMPOSITIONS

We, THE PROCTER & GAMBLE COMPANY, a Company organised under the laws of the State of Ohio, United States of America, of 301 East Sixth-Street, Cincinnati, Ohio 45202, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following

This invention relates to granular built laundry detergent compositions which provide simultaneous laundering and softening of textiles during conventional fabric faundering operations. Such compositions employ a combination of non-soap synthetic detergent compounds, organic or inorganic detergent builders and particular smextim

clay compounds having particular cution exchange characteristics.

Various clays materials have been used in many different types of detergent

without tayls mission are been used in analy untertax, yet o seedings systems for widely differing purposes. Cays, for example, have been disclosed for use so builders (Schwarze and Perry, Surjace Active Agents, Interscience Publishers, Inc., 1949, pp. 322 and 299); as water-softeness (British Petres 46), 221); as suri-ciking agents (U.S. Petres 2,625,733 and 2,779,607); as suspending agents (U.S. Petres 2,625,733); and as filter (U.S. Petres 2,625,738).

It is also well known that some clay materials can be deposited on fabrics to impart softening and antistatic properties thereto. Such clay deposition is generally realized by contacting fabrics to be so treated with aqueous clay suspensions (See,

for example, U.S. Patents 3,033,699 and 3,594,221).

Attempts, however, to incorporate clay materials into built detergent systems for the purpose of providing simultaneous fabric laundering and softening have not been entirely successful. Conventional detergent builders tend to retard or inhibit the tendency of clay materials to deposit on fabric surfaces, such deposition being necessary to realize the desired fabric softening results. Furthermore, to provide the requisite uniform deposition of clay material onto fabrics being laundered, the clay material must be thoroughly and quickly dispersed throughout the fabric laundering solution during the relatively brief wash cycle.

Some of these difficulties of providing through-the-wash clay softening have been resolved by using conventional fabric softening agents such as isostearic acid or polyamine or polyquaternary ammonium compounds in combination with clay in built detergent formulations (See U.S. Patents 3,594,212 and 3,625,905). The dispersability problem can be alleviated somewhat by adding to laundering solutions built liquid detergent compositions wherein clay is suspended and therefore more easily dispersed (See U.S. Patent 2,920,045). However, such liquid heavy-duty built laundry com-

positions do not provide the convenience associated with granular laundry products. Accordingly, it is an object of the present invention to provide granular built laundry detergent compositions which can yield simultaneous fabric laundering and

It has surprisingly been discovered that by using particular types of clay having particular cation exchange characteristics, these objectives can be realized and built

10

15

25

20

30

These phosphate esters can be modified by including in the molecule from one to about 40 alkylene oxide units, for example, ethylene oxide units. The formulae for these modified phosphate anionic detergrants are

55

15

20

25

30

40

45

50

10

15

20

30

35

45

50

in which R represents an alkyl group commining from about 8 to 20 curbon atoms, or an alkylphenyl group in which the alkyl group contains from about 8 to 20 carbon atoms, and M represents a soluble cution such as hydrogen, sodium, potassium, ammonium or substituted ammonium; and in which n is an integer from 1 to 40.

Another class of suitable anionic organic desergents particularly useful in this invention includes salts of 2-acyloxyalkane-1-sulphonic acids. The salts have the

where R<sub>1</sub> is alkyl having from 9 to 23 carbon atoms (forming with the two carbon atoms an alkane group); R2 is alkyl having from 1 to 8 carbon atoms; and M is a

water-soloble cation

The water-soluble carion, M. in the hereinbefore described structural formula can be, for example, an alkali metal cation (for example sodium, potassium, lithium), ammonium or substituted-animonium cation. Specific examples of substituted animonium cations include: methyl-, dimethyl-, and truncthyl- ammunium cations and quaternary ammonium cations such as tetramethyl-ammonium and dimethyl piperidinium cations and those derived from alkylamines such as onlylamine, dicthylamine, triothylamine and

and having the formula

Specific examples of beta-acyloxy-alkane-1-sulphonates, or alternatively 2-acyloxyalkane-1-sulphonates, meful herein include the audium salt of 2-acetoxy-tridecane-1sulphonic acid; the potassium salt of 2-propionyloxy-tetradecane-1-sulphonic acid; the lithium salt of 2-butanoyloxy-tetradecane-1-sulphonic acid; the solium salt of 2-tentanoyloxy-pentadecame-1-supposite acid; the sodium salt of 2-accessy-bessdecame-1-sulphonic acid; the possissium salt of 2-accessyloxyretradecame-1-sulphonic acid; the sodium salt of 2-acciony-heptadecane-1-sulphonic acid; the lithium salt of 2-acctonyoctadecane-I-sulphonic acid; the potassium salt of 2-acetoxy-nonadexnos-I-sulphonic acid: the sedium sait of 2-acetoxy-uncosane-1-sulphonic acid; the sedium sait of 2propionyloxy-docosane-1-sulphonic acid; the isomers thereof.

Preferred beth-fcyloxy-alkane-1-sulphonate saits herein are the alkali metal saits of beta-acetoxy-alkane-1-sulphonic acids corresponding to the above formula wherein R, is an alkyl having from 12 to 16 carbon atoms; these sales being preferred because

of their excellent cleaning properties and ready availability.

Typical examples of the above described beta-accounty alkane-sulphonaues are described in the literature: Belgium Patent 650,323 discloses the preparation of certain 2-acyloxy alkanesulphonic acids. Similarly, U.S. Patents 2,094,451 (Guenther et al) and 2,086,215 (DeGresse) disclose certain salts of beta-acroxy alkanesulphonic acids.

Another preferred class of anionic descripent compounds herein, both by virtue of superior cleaning properties and low sensitivity to water hardness (Ca++ and Mg++ ions) are the alkylated e-sulyphocarboxylates, containing from 10 to 23 carbon atoms

wherein R is C<sub>2</sub> to C<sub>20</sub> alkyl, M is a water-soluble cation as hereinbefore disclosed, preferably a sodium ions, and R is a C<sub>2</sub>—C<sub>2</sub> alkyl, e.g., methyl, ethyl, propyl and butyl. These compounds are prepared by the estertification of a-suphonanced carboxylic acids, which are commercially available, using standard techniques. Specific examples of the alkylated a-sulphocarboxylates preferred for use herein include:

Ammonium methyl-a-sulphopalmitate, Triethanolammonium ethyl-e-sulphostescare, sodium methyl-o-sulphopalmitate, sodium ethyl-e-sulphopolimitate,

35

40

10

15

20

25

30

35

40

50

sodium buryl-a-sulphostearate, potassium methyl-a-sulpholaurate. lithium methyl-a-sulpholaurate,

as well as mixtures thereof.

A preferred class of anionic organic detergents are the  $\beta$ -alkyloxy alkane sulphorsares. These compounds have the following formula:

where  $R_s$  is a straight chain alkyl group having from 6 to 20 carbon atoms,  $R_s$  is a lower alkyl group having from 1 (preferred) to 3 carbon atoms, and M is a water-10 soluble cation as hereinbefore described.

Specific examples of  $\beta$ -alkyloxy alkane-naphonanes, or alternatively 2-alkyloxy-alkane-1-sulphonates, having low hardness (calcium ion) sensitivity useful herein to provide superior cleaning levels under household washing conditions include:

potassium-8-methoxydecanesulphonate. 15 sodium 2-methoxytridecanesulphonate, potassium 2-ethocytetradecylsulphonate, sodium 2-isopropoxyhexadecylsulphonate, lithium 2-t-buttoxytetradecylsulphonare, sedium \$-merboxyoctadecylsulphonaue, and

20 ammonium B-a-propoxydodecylsulphonate.

Other synthetic anionic detergents useful berein are alkyl ether sulphates. These materials have the formula RO(C2H2O), SO2M wherein R is alkely or alkenyl having from 10 to 20 carbon atoms, x is from 1 to 30, and M is a water-soluble cation as defined hereinbefore. The alkyl other sulphaues useful in the present invention are 25 condensation products of ethylene oxide and monohydric alcohols having from 10 to 20

carbon stoms. Preferably, R has 12 to 18 carbon atoms. The alcohols can be derived from fats, for example, crement oil or ratiow, or can be synthetic. Lauryl alcohol and straight chain alcohols derived from tallow are preferred herein. Such alcohols are ocapia chain sussions curried iron allow or preserved ineria. Such accords are reacted with firm I to 30, and especially 6, mokar proportions of chiplene coulds and the resulting mixture of multicular species, having, for example, an average of 6 molecular species, having, for example, an average of 6 molecular species and neutralizate. Perfectly examples of alloyl either sulphates of the prosent inversion are: sodium cocoma talkyl chipme glycod cheb sulphate; lithium tallow alkyl triethylene glycod.

ether sulphate; and sodium tallow alkyl hexaoxyethylene sulphate.

Priesrod herein for reasons of excellent cleaning properties and ready availability are the alkali metal cocount- and tillow-skyl cayethylene ether sulphates having an average of from 1 to 10 oxyethylene modelies. The alky ether sulphates of the present invention are known compounds and are described in U.S. Patent 3,332,876 to Walker. Additional examples of anionic non-soap synthetic detergents which come within

the terms of the present invention are the reaction product of fatty acids esterified with isothionic acid and neutralized with sodium hydroxide where, for example, the fatty acids are derived from excount oil; sodium or potassium salts of fatty acid amides of methyl tauride in which the fatty acids, for example, are derived from cocount oil. Other anionic synthetic detergents of this variety are set forth in United States Patents 2,486,921; 2,486,922; and 2,396,278. 45

Additional examples of anionic, non-soap, synthetic detergence, which come within the terms of the present invention, are the compounds which commin two amonic functional groups. These are referred to as di-anionic devergents. Suitable di-anionic detergents are the disulphonates, disulphates, or mixtures thereof which may be represented

50 by the following formulae:

where R is an acyclic aliphatic hydrocarbyl group having 15 to 20 carbon atoms and M is a water-solubilizing cation, for example, the C, to C, disodium 1,2-alkyldisulphates,

5	1,400,898	- 5
	$C_{15}$ to $C_{26}$ dipotassium 1,2-alkyldisulphomates or disulphates, disodium 1,9-hexadecyl disulphates, $C_{12}$ to $C_{26}$ disodium-1,2-alkyldisulphomates, disodium 1,9-stearyldisulphates and 6,10-octadecyldisulphates.	
5 -	The aliphatic portion of the disulphates or disulphonates is generally substantially linear, thereby imparting desirable bin-degradable properties to the detergent compound.	5
	The water-solubilizing cations include the customary cations known in the detergent art, i.e., the alkali metals, and the ammonium cations, as well as other metals in group	
	HA, HB, HIA, IVA and IVB of the Periodic Table except for boron. The preferred	**
)	water-solubilizing extions are sodium or possssium. These dianionic detergents are more fully described in British Leners Pateur 1,151,392.  Still other autonic synthetic detergents include the class designated as succinamates.	-10
	This class includes such surface active agents as disodium N-octadecylsulphosuccina- mate; retrasodium N-(1,2-dicarboxyethyl)-N-octadecylsulpho-succinamate; diamyl ester	
5	of sodium sulphosuccinic acid; dihexyl ester of sodium sulphosuccinic acid; diocuyl esters of sodium sulphosuccinic acid. Other suitable anionic detergents utilizable herein are olefin sulphonates having	15
10	from 12 to 24 carbon atoms. The term "oleffia sulphonates" is used herein to mean compounds that can be produced by the sulphonation of so-folias by means of uncom- plexed sulphur trioxide, followed by neutralization of the soid neutron matters in con- ditions such that any sultones which have been formed in the reaction are hydrotyzed to give the corresponding hydroxy-alkame-sulphonates. The sulphur trioxide can be fleuid or gaseous, and is usually, but not necessarily, diluted by here dilutents, for	20
15	example by liquid SO <sub>2</sub> or chlorinated hydrocarbons, when used in the liquid form, for example by sir, nitragen or gaseous SO <sub>2</sub> , when used in the gaseous form.	2
,	The w-olefuls from which the oleful sulphonauts are derived are mone-olefuls having 12 to 24 carbon atums, preferably 14 to 16 carbon atums, Preferably, they are straight chain olefuls. Examples of suitable 1-olefuls include 1-dedecents: 1-terra-	
<b>30</b>	decree: 1-broadcerne: 1-octoderne: 1-citosene and 1-entracysen.  In addition to the true allenee sulphonases and a proportion of hydroxy-alkane- sulphonates, the olefin sulphonates can contain minor amounts of other materials, such as alkene disalphonates depending upon the reaction conditions, proportion of reactants, the nature of the starting olderins and impurities in the oldin stock and side reactions	3
35	during the sulphonation process.  A specific anionic detergent which has also been found excellent for use in the present invention is described more fully in the U.S. Patent 3,332,880 of Phillip F.	3
10	Pflaumer and Adrian Kessler.  Of all the above-described types of snionic surfactants, preferred compounds include sodium linear alkyl benzene sulphonace wherein the alkyl chain averages from about 10 to 18, more preferably about 12, carbon arous in length, sodium tallow alkyl	4
	sulphate, 2-activy-tridecane-1-sulphonic acid; sodium methyl-a-sulphopaliminare; sodium $\beta$ -methoxyoctadecylsulphonate; sodium occount alkyl ethylene glyxol ether sulphonate; the sodium sait of the sulphunic acid ester of the reaction product of one mole of fallow alcohol and three moles of ethylene oxide; and mixtures thereof.	
15	Ampholytic Synthetic Detergents.  Ampholytic Synthetic Detergents can be broadly described as derivatives of aliphatic or aliphatic derivatives of hencexcyclic secondary and territary amines in which the aliphatic radical may be straight chair or branched and wherein one of the aliphatic substituents contains from about 8 to 18 cuttors atoms and at least one contains and	4
60	anionic water-solubilizing group, for example, carboxy, sulphonane, sulphato. Examples of compounds falling within this definition are: sodium 3-(dode/caplamino)-propionate, sodium 3-(dode/slamino) propuse-1-sulphonare, sodium 2-(dodecylamino) ethyl sulphata; sodium 2-(disentifylamino)-confaceanone, disedium 3-(N-carboxymethyldo-	
55	decylamino)-propine-1-sulphonate, disodium octadecylimiaediazensie, sedium 1- carboxynethyl-2-undecylimidazode, and sodium N <sub>3</sub> N-bis(2-bydroxyethyl)-2-sulphatu- 3-dodecoxypropylamine. Sodium 3-(dodecylamino)propine: 1-sulphonate is preferred.	
	Zwitterionic Synthetic Detergents.  Zwitterionic surfactants can be broadly described as derivatives of secondary and	
0	nertiary amines, derivatives of heterocyclic secondary and tertiary amines, or derivatives of quaternary ammonium, quaternary phosphonium or tertiary sulphonium compounda. The cationic atom is the quaternary compound can be part of a heterocyclic ring. In all of these compounds there is at least one aliphatic group, straight chain or branched, constaining from 5 to 18 carbon atoms and as least one aliphatic substituent containing	

15

20

40

45

10

25

30

35

45

50

an anionic water-solubilizing group, for example, curboxy, sulphonate, sulphoto, phos-phato, or phosphono. Examples of various classes of zwitterionic surfactants operable herein are described as follows:

1. Compounds corresponding to the general formula

wherein R, is alkyl, alkenyl or a hydroxyalkyl containing from 8 to 18 carbon atoms and containing if desired up to about 10 ethylene oxide moieties and/or a glyceryl moiety, Y, is nitrogen, phosphorus or sulphur, R<sub>e</sub> is alkyl or monohydroxyalkyl containing 1 to 3 carbon atoms; x is 1 when Y2 is S, 2 when Y, is N or P; R2 is alkylene or hydroxyalkylene containing from 1 to 5 carton atoms; and Z is a carboxy, sulphonate,

sulphate, phosphotate group.

Examples of this class of zwitterionic surfactants include: 3-(N,N-dimethyl-N-di szampies of this class of zwiterionic surfactants include: 3-(N,N-dimethyls-N-hexadecylammonio)-propanel-taplipmant; 3-(N,N-dimethyls-N-beandecylammonio)-propanel-taplipmant; 3-(N,N-dimethyls-N-beandecylammonio)-this order of the surface of the s

propionate.

propocate.

Preferred compounds of this class from a connaercial standpoint are 3 - (N,N-dimethyl - N - hexadecylamanoslo) - 2 - hydroxypropene - 1 - sniphocane; 3 - (N,N-dimethyl - N - allykamnosno) - 2 - hydroxypropene - 1 - sniphocane; 3 - (N,N-dimethyl - N - hexadecylamnosol) - propulse.

- subjective i from tallow fatry alcohol; 3 - (N,N - dimethyl - N - hexadecylamnosol) - propulse.

- subjective i from tallow fatry alcohol; 3 - (N,N - dimethyl - N - trendecylamnosol) propune - 1 - sniphocane; 2 - (N,N - dimethyl - N - stendecylamnosol) - 2 - hydroxypropune - 1 - sniphocane; the sniph group being derived from the middle ent of eccount fany alcohol; adjobnost; the sniph group being derived from the middle ent of eccount fany alcohol; 25

sulphonate, the alkyl group being derived from the minute cut or covonate lawy massars, 3 - (NN) - dimethyl-decephramsonis) - 2 - hydroxygropane - 1 - sulphonate; <math>4 - (NN) - dimethyl - terradecylammonis) houther - 1 - sulphonate; <math>4 - (NN) - dimethyl - terradecylammonis) beamse - 1 - sulphonate; <math>4 - (NN) - dimethyl - hendecylammonis) beamse - 1 - sulphonate; <math>4 - (NN) - dimethyl - hendecylammonis) beamse - 1, <math>2 - (NN) - dimethyl - N - cctaclecylammonis) beamse - 1, <math>2 - (NN) - dimethyl - N - cctaclecylammonis) beamse - 1, <math>2 - (NN) - dimethyl - N - cctaclecylammonis) - 2 - methyl cyponer - 1 - sulphonate; <math>3 - (NN) - dimethyl - N - cctaclecylammonis)35

dimenty - N - encosystemators of the second German Patent L018.421.

2. Compounds having the general formula:

wherein R, is an alkyl, cycloalkyl, aryl, amikyl or alkaryl group containing from 10 to 20 carbon atoms; M is a bivulent radical selected from: aminocarbonyl, carbonylamino, carbonyloxy, aminocarbonylamino, the corresponding this groupings and substituted amino derivatives;  $R_s$  and  $R_s$  are alkylene groups containing from 1 to 12 carbon atoms;  $R_{\phi}$  is alkyl or hydroxyalkyl containing from 1 to 10 carbon atoms;  $R_{\phi}$  is selected from  $R_{\phi}$  groups,  $R_{\phi} = M_{\phi} - M_{\phi}$ ; and  $-R_{\phi}$  COOMe wherein  $R_{\phi}$   $R_{\phi}$   $R_{\phi}$  and  $R_{\phi}$  are as defined above and Me is a monovalent salf-forming cathon. Components of this

30

35

10

15

20

30

pared in accordance with methods described in U.S. Patent 3,265,719 and German Auslegerschrift (Published Specification) 1,018,421 3. Compounds having the general formula:

wherein  $R_{\rm u}$  is an alkyl group,  $R_{\rm zo}$  is a hydrogen atom or an alkyl group, the total number of carbon atoms in  $R_{\rm u}$  and  $R_{\rm ze}$  being from 8 to 16 and

represents a quaternary annuous group in which each group  $R_{\rm NL}$   $R_{\rm NL}$  and  $R_{\rm 10}$  is an airyl or hydroxyalityl group or the groups  $R_{\rm NL}$   $R_{\rm NL}$  and  $R_{\rm NL}$  are consoland in a heterocyclic ring and n is 1 or 2. Enempties of satisfacts extitexions; variations of this betterocyclic rine and n is 1 or 2. Enempties of satisfact extitexions arisinfractures of the include the  $\gamma$  and  $\delta$  hexadecyl pyrinion sulphobectsines, the  $\gamma$  and  $\delta$  hexadecyl  $\gamma$ -prioritor salphobectsines, the  $\gamma$  and  $\delta$  the contradiction of such a system of the contradiction of such a system of the contradiction of such a system of the form of the published South African permit applications of  $\delta$  4,5788.

4. Compounds having the general formula 10 15

wherein R<sub>ss</sub> is an alkarylmethylene group containing from 8 to 24 carbon atoms in the alkarylmin N<sub>ss</sub> is selected from R<sub>ss</sub> groups and alkyl and hydrosynkly groups contain-ing from 1 to 7 orthon atoms, R<sub>ss</sub> is alkyl to hydrosynkly, containing from 7 orthon atoms, R<sub>ss</sub> is alkylene or hydrosynklytene containing from 1 to 7 carbon atoms and the second from the 20 25

and Z<sub>1</sub> is selected from: sulphomairs, carbony and sulphairs. Rampiles of a switter-food surfaceants of this type include 3 - N. - dodes/sbenzyl - N.N. - dimethylammonis)-propane - 1 - sulphomairs; 4 - (N - dodes/sbenzyl - N.N. - dimethylammonis)-propane - 1- sulphomairs; 5 - (N - dodes/sbenzyl - N.N. - dimethylammonis)-propanes; 1 - sulphomairs; 5 - (N - dodes/sbenzyl - N.N. - dimethylammonis)-propinents; 4 - (N - dodes/sbenzyl - N.N. - dimethylammonis)-propinents; 4 - (N - dodes/sbenzyl - N.N. - dimethylammonis)-propinents; 4 - (N - dodes/sbenzyl - N.N. - dimethylammonis)-propinents; 5 - (N - dodes/sbenzyl - N.N. - dimethylammonis)- n. - dodes/sbenzyl - N.N. - dimethylammonis)- n. - dodes/sbenzyl - N.N. - dimethylammonis)- n. - dodes/sbenzyl - N.N. - dimethylammonis - N.N. - dimethylammonis - 1 - (N - dodes/sbenzyl - N - methylammonis)- n. - dodes-sbenzyl - N - methylammonis - n. - dodes-sbenzyl - N - methylammonis - n. - sulphomair - n. - differents-sbenzyl - N - methylammonis - n. - sulphomair - n. - differents-sbenzyl - N - methylammonis 2-bydroxypropane-1-salphonate.

Zwitterionic surfaciants of this type as well as methods for their preparation are described in U.S. Patents 2,697,116; 2,697,656 and 2,669,991 and Canadian Patent 883,864.

5. Compounds having the general formula:

wherein R<sub>10</sub> is an alkylphenyl, cycloalkylphenyl or alkenylphenyl group containing from 8 to 20 carbon aroms, in the alkyl, cyclosilkyl or alkenyl moiety;  $R_{1a}$  and  $R_{2a}$  are each aliphatic groups containing from 1 to 5 carbon atoms;  $R_{2a}$  and  $R_{2a}$  are each hydrogen

	atoms, hydroxyl groups or aliphatic groups containing from 1 to 3 carbon atoms and $R_{\rm ex}$ is an alkylene group containing from 2 to 4 carbon atoms. Examples of zwitteriousic surfaceants of this type include 3-{N-dodecylphenyl-	
5	N.N dimethylammonio)propane - 1 - sulphonate; 3 - (N - hexadecylphenyl - N.N dimethyl)butane - 1 - sulphonate; 3 - (N - tetradecylphenyl - N.N dimethyl-ammonio) - 3.3 - dimethylmonane - 1 - sulphonate and 3 - (N - dodecylphenyl - N.N dimethylmonane - 1 - sulphonate and 3 - (N - dodecylphenyl - N.N dodecylphenyl - dodecylphenyl - N.N dodecylphen	5
	dimethylammanis) - 3 - hydroxypropane - 1 - sufptionare. Compounds of this type are described move fully in British Patents 970,883 and 1,046,252. Of all the above-described types of zwittenionic surfactants, preferred compounds	•
10	mchide: 3(N,N - dimethyl - N - alkylammonio) - propone - 1 - sulphonate and 3- (N,N - dimethyl - N - alkylammonio) - 2 - hydroxypropane: 1 - sulphonate wherein in both compounds the alkyl group averages 14.8 carbon aroms in length; 3(N,N - dimethyl - N - hexadecylammonio) - propane: 1 - sulphonates; 3(N,N - dimethyl-	10
15	N - bexadecylammonio) - 2 - hydruxyjrospane - 1 - sulphonane; 3 - (N - dedezyl-benzyl - NN - dimethylammonio) - propane - 1 - sulphonate; (N - dedezylbenzyl-NN - dimethylammonio)acetane; 3 - (N - dadecylbenzyl-NN - dimethylammonio)acetane; 3 - (N - dadecylbenzyl-NN - dimethylammonio)acetane; and (N,N-dimethylammonio)acetane.	15
	Builder Salts	
20	The detergent compositions of the present invention contain, as an essential com- ponent, a polyamionic detergent builder salt. In the present compositions these water- soluble builder salts serve to maintain the pH of the laundry solution in the range of from 7 to 12, preferably from 8 to 11. Furthermore, these builder salts enhance the	20
25	fabric cleaning performance of the overall compositions while at the same time serve to suspend particulate soil ecleaned from the surface of the fabrics and prevent its codeposition on the fabric surfaces. Surprisingly, although the detergency builder sales serve to suspend day soils of the kaolinite and illite types and prevent their redeposi- tion on fabrics, they do not appear to interfere with the deposition on fabric surfaces.	25
30	of the succtive-type clays softeners used herein. Furthermore, these polyandaric builder saits have been found to cause the senective-type clays present in the granular detergent formulations of the invention to be readily and homogeneously dispersed throughout the aqueous fundering medium with a minimum of agintion. The homogeneity of the clay dispersion is necessary for the clay of function effectively as a fabric softener,	30
35	while the ready dispersability allows generaler detergent compositions to be formulated. Suitable detergent builder salts useful herein can be of the poly-valent inorganic and poly-valent organic types, or mixtures thereof. Non-limiting examples of suitable water-soluble, inorganic detergent builder salts include the alkali metal carbonates, horates: Invastuates, nodemberatures trivilenthosophores, biographeses and existence and existence.	35
40	Specific examples of such salts include the sodium and porassium retrasborates, per- borates, bicarbonates, carbonates, tripolyphosphares, orthophosphares and hexameta- phosphare.	40
45	Examples of suitable organic altains detergency builder sits are: (1) water- soluble amino polyacetates, for example, sodium and potassim ethylenedismine terra- acetates, nitrilorincetates and N-(2-hydroxyethyl)neritodiacetates; (2) water-sobible saits of phyric acid, for camples, sodium and potassim physics; (3) water-sobible polyphosphusates, including, sodium, potassima and lithium sits of chance-1-hydroxy- ll-diptosphonic acid; sodium, potassima and lithium sits of enthelpendiphosphonic	<sup>-</sup> 45
	acid.  Additional organic builder salts useful herein include the polycarboxylare manirials	
50	described in U.S. Paraez 2,264,103, including the water-soluble albili metal salts of mellitic acid. The water-soluble salts of polycarboxylate polyumes and copolymess such as are described in U.S. Paraezt 3,308,067, are also smitable for use herein. It is to be understood that while the albali metal salts of the foregoing incorpanic and organic poly-	50
55	valent anionic builder salts are preferred for use herein from an economic standpoint, the ammentum, alkanolammonium, for example triethanolammonium ad diethanol- ammonium, water-soluble salts of any of the foregoing builder anions are useful	55
60	herein.  Mixtures of organic and/or inorganic builders can be used herein. One such unitarue of builders is disclosed in Canadian Patent 755,138, for example, a ternary-instrume of sodium tripolyphosphate, trisodium nitrilotriacente and trisodium ethano-laydrony-la-diphosphanate.	60
	While any of the foregoing alkaline poly-anionic builder materials are useful berein, sodium tripolyphosphate, sodium mitrilotriacetate, sodium mellitate, sodium	

S

10

20

25

30

35

45

50

60

S

10

15

20

25

20

35

50

55

60

citrate and sodium carbonate are preferred herein for this builder use. Sodium tripolyphosphate is especially preferred herein as a builder both because of its desergency builder activity and its ability to disperse homogeneously and quickly the smeetite clays throughout the aqueous laundry media without inverfering with clay deposition on the fabric surface. Sodium tripolyphosphate is also especially effective for suspending illite and knolinite clay soils and retarding their redeposition on the fabric surface.

The detergent builders are used at concentrations of from 10% to 60%, preferably from 20% to 50% by weight of the detergent compositions of this invention.

Clay Compounds.

The third essential component of the present compositions consists of particular smootite clay materials to provide fabric softening concurrently with fabric cleansing. These smeetite clays are present in the detergent compositions in amounts from 1% to 50%, preferably from 5% to 15% by weight of the total compositions. The clay minerals used to provide the softening properties of the instant composi-

tions can be described as expandable, three-layer clays i.e., alumino-silicates and magnesium silicates, having an ion exchange capacity of at least 50 may/100 g. de clay. The term "expandable" as used to describe clays relates to the ability of the layered clay structure to be swollen or expanded, on contact with water. The threelayer expandable clays used herein are those materials classified geologically as

smeetites.

There are two distinct classes of smectite-type clays: in the first, aluminium oxide is present in the silicate crystal lattice; in the second class of smectices, magnesium oxide is present in the silicate crystal lattice. The general formulae of these smecrites are  $A_a(Si_aO_a)_a(OH)_a$  and  $Mg_a(Si_aO_a)_a(OH)_{2a}$  for the aluminium and magnesium oxide type day, respectively. It is to be recognised that the range of the water or hydration in the above formulas can vary with the processing to which the day has been subjected. This is immaterial to the use of the smectire clave in the present invention in that the expandable characteristics of the hydrated days are dictated by the silicate lattice structure. Furthermore, arom substitution by iron and magnesium can occur within the crystal lattice of the smectices, while metal cations such as Na+, Ca++, as well as H+, can be co-present in the water of hydration to provide electrical neutral. ity. Except as mentioned below, such cation substitutions are immaterial to the use of the clays herein since the desirable physical properties of the clays are not sub-

stantially altered thereby. The three-layer, expandable alamino-silicates useful herein are further characterized by a dioctahedral crystal lattice, while the expandable three-layer magnesium

ailicates have a trioctahedral crystal lattice.

As noted hereinshove, the clays employed in the compositions of this invention contain cationic counterions such as protons, sodium ions, potassium ions, calcium ions and magnesium ions. It is customary to distinguish between clays on the basis of one cation predominantly or exclusively absorbed. For example, a sodium clay is one in which the absorbed cation is predominantly sodium. Such absorbed cations can become involved in exchange reactions with cations present in aqueous solutions. A typical exchange reaction involving a spectite-type clay is expressed by the following 45 equation:

## smectite day (Na)+NH,OHe≥smectité day (NH,)+NaOH

Since the foregoing equilibrium reaction, one equivalent weight of ammonium ion replaces an equivalent weight of sodium, it is customary to measure exchange capacity reguects are equarement weights or sourcing, it is consonant to measure extensing capacity (sementimes termined "base exchange capacity") in terms of milliogravitations per 100 g, of clay (meag-/100gs.). The curion exchange capacity of clays can be measured in several ways, including by electrodialysis, by exchange with ammonistum for followed by titration or by a methylene blue procedure, all as fully set forth in Grimshaw, "The Chemistry and Physics of Clays", pp. 264—265, Interestione (1971). The cation exchange capacity of a clay mineral relatus to such factors as the expandiable proverties of the elex, the charge of the class, which in turns, is determined at least in properties of the clay, the charge of the clay, which, in turn, is determined at least in part by the lattice structure. The ion exchange capacity of clays varies widely in the range from about 2 meq/100 g, for kaolinites to about 150 meq/100 g., and greater,

for certain clays of the montmorillionite variety. Illite clays have an ion exchange capacity somewhere in the lower portion of the range, i.e., about 26 meq/100 g. for an average illite clay. It has been determined that illite and kaolinite clays, with their relatively low ion exchange capacities, are not useful in the instant compositions. Indeed, such illine and

10	1,40x,898	10
5	kaolinite clays constitute a major component of clay soils and, as noted above, are removed from fabric surfaces by means of the instant compositions. However, smexities, such as nontrontle, having an ion exchange capacity of approximately 50 mag/100 gs, saponite, which has an ion exchange capacity of around 70 men/100 gs, and mentmoril- lonite, which has an ion exchange capacity of around 70 men/100 gs. have	5
	lonite, which has an ion exclusing capacity greater than 70 meo/100 g, have been found to be useful in the compositions of the invention in that they are deposited on the fabrics to provide the desired softening benefits. Accordingly, clay minerals useful herein can be characterized as expandable, three-layer snectine-type clays having an ion exchange capacity of at least about 50 mea/100 g.	
10	While not intending to be limited by theory, it appears that the advantageous softening (and potentially due sourceignig), benefits of the instant compositions are ascribable to the physical characteristics and ion exchange properties of the clays used therein. That is to say, experiments have shown that non-expandable clays such as the kardinities and the illitties, which are both classes of clays having four exchange	10
15	capacities below 50 may/100 g., do not provide the beneficial aspects of the days employed in the present compositions. Furthermore, the unique physical and electro- chemical properties of the smeetite days apparently cause their interaction with, and disporation by, the poly-amionic builder salts used in the present compositions. Thus,	15
20	it has now been found that, rather than agglomenting to form viscous gels when contacted by water, the smeetite days used herein can be added to aqueous laundry boths in granular compositions containing poly-anionic detergency builders of the type dischord herein ny yield homogeneous, stable day suspensions. The problems of gelling and agglomention usually encountered when smeetite clays are added to aqueous media in solid form are allewind by the presence of the builder. Apparently, the	20
25	negative electrical charges on the builder anions serve to repuise the clay particles, thereby providing the desired homogeneous day dispension and preventing agglomera- tion. Whatever the reason for the advantageous co-exting of the decregacy builder and succettle clays used herein, the combination of the poly-unionic detergency builders with the expendable, three-layer, discarbeiral alumino-silvers and removable, three-layer.	25
30	layer, triocalabedral magnesium silicares provides a means whereby such emectite days can be added to a sufficient-containing media in solid form to provide the homo- geneous clay dispersion required for effective febric softening.  The smocture clays used in the compositions herein are all commercially available. Such clays include, for example, montimorillonite, yelchorskorite, necurswite, hesturine,	30
35	supposite and suscenite. The clays berein are available under various tradenances, for example, Thirogel No. 1 and Gelwhite GP from Georgia Kasalis A., Ellizabeth, New Jersey; Volckiy BC and Volckiy No. 325, from American Colloid Co., Stockie, Illinois; Black Hills Bactorite BH459, from International Minerals and Chemicals; and Vergum Pro and Vergum F, from R. T., Vanderbilt, It is no be recognized that such susceins-	35
40	type minerals obtained under the foregoing tradenances can comprise minerals of the various discrete mineral entities. Such mixtures of the smecrite minerals are suitable for use herein.  While any of the smectite-type clays having a cation exchange caracity of at least	40
45	about 50 mea/100 g. are useful herein, certain days are preferred. For example, Gel- white GP is an extremely white form of smectize clay and is therefore preferred when formulating white granular desceptor compositions. Volclay BC, which is a smecting- type clay maineral containing as fests 3% of iron (expressed as Fe,O.) in the crystal- lattice, and which has a very high into exchange capacity, is one of the most efficient and effective clays for use in laundry compositions and is perferred from the standpoint	45
50	of product performance. On the other hand, certain succities clays marketed under the name "bentonin" are sufficiently contaminated by other silicate meta-sits that their ion exchange capacity falls below the requisite range, and such clays are of no use in the compositions of this invention.  Appropriate clay minerals for use herein can be selected by virtue of the fact that	50
55	smeatities exhibit a true 14 a x-ray diffraction pattern. This characteristic pattern, taken in combination with exchange capacity measurements performed in the manner mored above, provides a basis for selecting particular smeatite-type minerals for use in the granular detergent compositions disclosed herein.	55
60	Optional Components.  The detergent compositions disclosed berein can contain other materials commonly used in such compositions. For grounds, we can all present the compositions are contained to the composition of the composition of the components.	60

The detergent compositions disclosed herein can constain other materials commonly used in such compositions. For example, various sull-suspending agents each as earbeonmetaly-callulose, corrosion inhibitors, duye, fillers such as codium sulphate and silica, optical brightness, such sources, such depressants, genricides, such-tamishing agents, pH adjusting agents such as sodium silicate, cueyanes, and the like, well-kinowi in the

11	1,400,898	11
5	art for use in detergent compositions, can also be employed herein. Bound water can also be present in said detergent compositions.  The clay-containing detergent compositions of this invention are in granular form. The compositions can be prepared by simply admixing the appropriate aignredients in the young the compositions can be prepared by simply admixing the appropriate aignredients in they form. The compositions are then added to water to provide a laundering liquor containing the instant compositions to the extent of from 0.02% to 2% by weight. Solled fobtics are added to the laundering liquor and cleansed in the usual memory. The	5
	effective amount of the descriptor compositions to be used will depend to an extent on the weight of clothes being laundered and their degree of soiling. Aqueous laundering	
10	haths containing said compositions provide adequate clearing and softening benefits with solicid fabrics, especially control and conton/polyeter bleachs. The suspended day material found in the laundering liquor also serves to absorb fugitive dye in solution, thereby reducing or inhibition dwe transfer.	10
15	The granular built detargent compositions and the fabric handering and soften- ing process of the present invention are illustrated by the following examples. Desired cotton terry washedotts were washed in aqueous solutions having dissolved thenche various day-cuttaining built granular detergent compositions of this invention. Soft- ness of the terry swatches so washed was compared with the softness of terry wastrhes	15
20	washed in an equivalent concentration of the same built granular detergent without the clay, as well as with the softness of earry swatches washed in this same no-clay deter- gent solutions followed by rinsing to water containing a commercially available fabric softners, Downy. Composition and solution concentrations are described in Table I below.	20
25	The terry swatches were washed for 10 minutes in a minuture agitator containing two gallons of washing liquor at 120° F. and 7 gr/gel, artificial hardness. The swatches comprised 4%, by weight of the washing liquor. After washing, the swatches were spun dry and rinsed with two gallons of water at 120° F. and 7 grains/gallon artificial hardness. Swatches were then dried in a conventional electric driver.	25
30	After several treatment cycles, the test and control swatches were graded taccilely for softness by a pasted of three to five judges making paired comparisons of all swatches. Graders assigned an integer grade of from 0 to 4 on a linear scale to the softer treatment of each pair, assigning the higher grades to coverspending larger differences in softness. The data considered were analyzed statistically no obtain mean	30
35	softness grades (panel score units) for each treatment and a statistical estimate of the least significant difference (LSD) at the 93% confidence level. Results of the softening tests appear in Table I.	35

۲

10

15

TABLE I

		Composition No.					
Component-Wt. %	1	2	3	4	5	6	
Anionic Surfactant	* 16.8	16.8	16.8	15.3	8.4	16.8	
Sodium tripely- phosphate	32.9	32.9	32.9	45.0	24.7	49.5	
Sodium Silicate	5.9	5.9	5.9	5.37	2.9	5.9	
Sodium Sulphate	19.6	29.6	29.6	12.8	7.0	14.t	
Miscellancons minors Gelwhite GP**	~4.1 I0.0	~4.1	~4.3	<b>~</b> 2.8	~1.6	3.1سم	
Voiciny BC***				9.1	50.0		
Moisture	Balance	Balanco	Balance	Balance	Balance	Balance	
Solution Concen- tration (wt %) of Composition	0.104	0.104	0.104	0.11	0.20	0.10	
Solution pH	9.2	9.2	9.2	9.3	9.3	9.2	
Rinse	Water	Water	Downy (0.07% wt.)	Water	Water	Water	
Number of Cycles	4	4	4	2	2	2	
Mean Softness Grade (Panel Score Units)	0.8	-2.1	0.2	-0.5	1.7	-2.6	
Least Significant Difference (LSD	)	0.9			1.0		

\* A mixture in a 1,22:1 wt. ratio of sodium tallow sikel sulphate and sodium linear alkyl benzene sulphonate wherein the alkyl chain of the sulphonate averages 11.8 carbon atoms in length.

\*\* A commercially-available sodium montmorillonite clay having an ion-exchange capacity of about 100 meq./100 g.

\*\*\* A commercially-available sodium montmorillonite clay having an ion-exchange capacity of about 85-100 meg./100 g.

It can be seen from Table I that Compositions 1, 4 and 5 of the present invention provide softening benefits superior to built-detergent formulations commining no day softening agents and softening benefits compamble to those obtained with a commercial fabric softening rinse additive.

Compositions 1, 4 and 5 of the present invention also provide excellent cleaning

and detergency when employed in washing solutions at the specified concentrations, bubstantially similar detergency and solutions prosults were obtained when the amount surfactant mixture in Composition 1, 4 or 5 (Table 1) was replaced with an equivalent amount of 2-acrtoxy-tridecame-1-sulphonic acid; sodium methyl-s-sulphonpalmitate; sodium  $\beta$ -methoxyoczadecylsulphonate; sodium cucount alkyl ethylene glycol ether sulphonate or the sodium salt of the sulphuric acid ester of the reaction product of one mole of tallow fatty alcohol and three moles of ethylene oxide.

Substantially similar denergency and softening were obtained when the anionic surfactant mixture in Composition 1, 4 or 5 (Table I) was replaced by an equivalent 15

13	1,400,898				
\$	amount of 3 (N.Ndimethyl-N-altylammunia)- propase-1-sulphanaet or 3 (N.Ndimethyl-N-altylammunia)-2 hapknyamtenylammunia)-2 hapknyamtenylammunia)-2 hapknyamtenylammunia)-2 hapknyamteylammunia)-2 hapknyamtey 3 (N.Ndimethyl-N-bacadecylammunia)- propase -1 - sulphonate; 3 (N.Ndimethyl-N-bacadecylammunia)-2 - hvolroxypropase -1 - sulphonate; 3 - (N dodecylbezyl - N.N dimethylammonia)-anominia)- propase -1 - sulphonate; (N dodecylbezyl - N.N dimethylammonia)-acetase; 3 - (N dodecylbezyl - N.N dimethylammonia)-acetase; 3 - (N dodecylbezyl - N.N dimethylammonia)-2 - N.N dimethylammonia (N.N dimethylammonia)-3 - N.N dimethylammonia (N.N dimethylammoni	5			
10	decylammonio)-acetare, or sodium 3 - (dodecylamino)pixoyane - 1 - sulphunate. Substantialty similar detergency and softening were obtained when the sodium tripolyphosphate builder in Composition 1, 4 or 5 ('Table I) was replaced by an equivalent amount of sodium nitrilorriacetate, sodium mellitate, sodium citrate or sodium carbonate.	10			
15	Substantially similar detergency and sofrening were obtained when the clay softening agent in Compositions 1, 4 or 5 (Table 5) was replaced by an equivalent amount of volchonshoite, nontronite, herorotic or sauconite, all such clays having an ins-exchange capacity greater than 50 meq./100 g.  In addition to the unexpected their softening benefits which the built laundry	15			
20	detergent compositions of this invention provide, there are other advantages which this invention makes possible. For instance, dye-transfer inhibition, noted above, is a significant advantage not commonly shared by ordinary fabric selfening compositions. Moreover, the particular class of clays described herein which are deposited on the fabrics, provide a solv-fease benefit. The clays are adsorbed by the fabrics being better the class are adsorbed by the fabrics being the class of the class are advantaged by the class are advantaged by the fabrics being the class of the class are advantaged by the fabrics being the class of the class are advantaged by the fabrics being the class of the class are advantaged by the class	20			
25	wasted providing an improved soil-release surface. The benefit from this treatment is that during subsequent washing, suitan and soils are more easily removed from the fabrics in comparison with a fabric which has not previously been exposed to a treatment by the clay-containing compositions of this inventions. Still further, all of these benefits are enjoyed without impairing the water-sharpent malifies of the treated fabric. This	25			
30	is in marked contrast with ordinary quaternary arminonium fabric softmers which may tend to reduce the water-absorbent property of treated fabrics after several cycles. It is especially significant that each of the advantages described above in no way impairs or unterferes with the general overall cleaning effectiveness of the detection composition. The fact that these achievements are attained during the relatively brief span of a short washing cycle, for example from 6 to 15 minutes, is expectally noto-	30			
20	worthy.  WHAT WE CLAIM IS:—  1. A granular, built handry detergent composition comprising:	35			
40	<ul> <li>(a) from 2% to 30% by weight of a non-wap synthetic detergent compound selected from: anionic synthetic detergents, amphotytic synthetic detergents, zwitterhouse synthetic detergents and mixtures thereof;</li> <li>(b) from 10% to 60% by weight of one or more organic and/or inorquaic detergent builder safts; and</li> <li>(c) from 1½, to 50% by weight of a fabric-softening agent that is an expandable, three-layer smeetise-type day having an ion exchange capacity of at less 50</li> </ul>	40			
45	meq/100 g, said composition providing a solution pH of from 7 to 12 when dissolved in water at a concentration of 0.12%, by weight. 2. A composition according to claim 1 which contains as component (a) from 5-20%, by weight of one or more synthetic detergent compounds.	45			
50	3. A composition accreting to claim 2, wherein as emponent (a) one or more anionic synthetic detergent compounds are present. 4. A composition according to any one of claims 1—3, wherein the anionic deergent is a water soluble organic sulphote or sulphonate containing an alkyl group having 8 to 22 carbon atoms.	50			
S5	5. A composition according to any one of claims 1—3, wherein the anionic detergent is selected from solding linear alkyl benzene sulphorate having an average of 10 to 18 carbon arouns in the alkyl group, sediam fallow alkyl sulphane, 3-acctive/printecano-1-sulphonic acid, sodium methyl—sulphopalmitate, sodium β-methoxy octabecyl sulphonate, sodium coconat alkyl ethylene gived their authorate the sodium satisfactors.	55			
60	sulphuric acid ester of the reaction product of one mol of tallow fatty alcohol and three moles of ethylene oxide, and mixtures thereof.	60			

20 to 50% by weight of one or more originic and/or inorganic descrepct builder salts.
8. A composition according to any one of claims 1—7, wherein component (b) is selected from: alkali metal carbiovates, bicarbouates, borates, phosphates, polyabosphates, sulphates; water soluble antinopolyacentes, physians and poly phosphosates.
9. A composition according to any not of claims 1—7 wherein component (b) is

 A composition according to any one of claims 1.—7 wherein component (b) is selected from sodium tripolyphosphate, sodium nitrilotriacetane, sodium melitrare, sodium cirate and sodium carbosate.

10. A composition according to any one of claims 1—9, which contains from 5 to 15% by weight of the smectite-type clay.

10

15

20

15 15%, by weight in its smeather-type casy.
15 11. A composition according to any one of claims 1—10 wherein the smeatite-type clay is selected from montmorillouities, volchooskoites, noutronites, hectorites, saponites and sauconites.

12. A composition according to claim 1 substantially as described in the Examples

20 13. A process for the simultaneous laundering and softening of fabrics, comprising contacting said fabrics with an aqueous medium containing from 9.02% by weight of a composition in accordance with claims 1—12.

For the Applicants, CARPMAHLS & RANSFORD, Chartered Patent Agents, 43, Bloomsbury Square, London, WC1A 2RA.

Printed for Her Majesty's Stationery Office by the Courier Press, Learnington Spa, 1875. Published by the Patent Office, 25 Southampton Enddings, London, WC2A IAY, from which captes may be obtained.